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(54) Title: METHODS FOR STOOL SAMPLE PREPARATION

(57) Abstract

The present invention provides methods for the preparation of stool samples to increase the yield of relevant DNA, and further provides methods for isolating and analyzing target DNA for characteristics indicative of colorectal cancer. Methods for screening patients for the presence of cancerous or pre-cancerous colorectal lesions are also provided.

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METHODS FOR STOOL SAMPLE PREPARATION

FIELD OF THE INVENTION

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This invention relates to methods for the early detection of colon cancer in patients, and more particularly to methods for preparing stool samples in order to increase the yield of nucleic acids.

BACKGROUND OF THE INVENTION

Stool samples frequently must be prepared for medical diagnostic analysis. Stool samples may be analyzed for diagnosis of medical conditions ranging from parasitic, bacterial or viral infections to inflammatory bowel disease and colorectal cancer.

Colorectal cancer is a leading cause of death in Western society. However, if diagnosed early, it may be treated effectively by removal of the cancerous tissue. Colorectal cancers originate in the colorectal epithelium and typically are not extensively vascularized (and therefore not invasive) during the early stages of development. Colorectal cancer is thought to result from the clonal expansion of a single mutant cell in the epithelial lining of the colon or rectum. The transition to a highly vascularized, invasive and ultimately metastatic cancer which spreads throughout the body commonly takes ten years or longer. If the cancer is detected prior to invasion, surgical removal of the cancerous tissue is an effective cure. However, colorectal cancer is often detected only upon manifestation of clinical symptoms, such as pain and black tarry stool. Generally, such symptoms are present only when the disease is well established, and often after metastasis has occurred. Early detection of colorectal cancer therefore is important in order to significantly reduce its morbidity.

Invasive diagnostic methods such as endoscopic examination allow for direct visual identification, removal, and biopsy of potentially cancerous growths. Endoscopy is expensive, uncomfortable, inherently risky, and therefore not a practical tool for screening populations to identify those with colorectal cancer. Non-invasive analysis of stool samples for characteristics indicative of the presence of colorectal cancer or

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precancer is a preferred alternative for early diagnosis, but no known diagnostic method is available which reliably achieves this goal.

Current non-invasive screening methods involve assaying stool samples for the presence of fecal occult blood or for elevated levels of carcinoembryonic antigen, both of which are suggestive of the presence of colorectal cancer. Additionally, recent developments in molecular biology provide methods of great potential for detecting the presence of a range of DNA mutations or alterations indicative of colorectal cancer. The presence of such mutations can be detected in DNA found in stool samples during various stages of colorectal cancer. However, stool comprises cells and cellular debris from the patient, from microorganisms, and from food, resulting in a heterogeneous population of cells. This makes detection of small, specific subpopulations difficult to detect reliably.

An additional problem encountered in preparation of stool sample for detection of colorectal cancer is the difficulty of extracting sufficient quantities of relevant DNA from the stool. Stool samples routinely contain cell debris, enzymes, bacteria (and associated nucleic acids), and various other compounds that can interfere with traditional DNA extraction procedures and reduce DNA yield. Furthermore, DNA in stool often appears digested or partially digested, which can reduce the efficiency of extraction methods.

20 SUMMARY OF THE INVENTION

It has now been appreciated that the yield of nucleic acid from a stool sample is increased by providing an optimal ratio of solvent volume to stool mass in the sample. Accordingly, the invention provides stool sample preparation protocols for increasing sample nucleic acid yield.

In a preferred embodiment, methods of the invention comprise homogenizing a representative stool sample in a solvent in order to form a homogenized sample mixture having a solvent volume to stool mass ratio of at least 5:1. DNA may then extracted from the homogenized sample mixture. Providing an optimal solvent volume to stool mass ratio increases the yield of nucleic acid obtained from the sample. An especially-preferred ratio of solvent volume to stool mass is between about 10:1 and about 30:1, more preferably from about 10:1 to about 20:1, and most preferably 10:1.

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A preferred solvent for preparing stool samples according to the invention is a physiologically-compatible buffer, such as a buffer comprising Tris-EDTA-NaCl. A preferred buffer is a Tris-EDTA-NaCl buffer comprising about 50 to about 100 mM Tris, about 10 to about 20 mM EDTA, and about 5 to about 15 mM NaCl at about pH 9.0. A particularly preferred buffer is 50 mM Tris, 16 mM EDTA and 10 mM NaCl at pH 9.0. Another preferred solvent is guanidine isothiocyanate (GITC). A preferred GITC buffer has a concentration of about 1 M to about 5 M. A particularly preferred GITC buffer has a concentration of about 3 M.

Also in a preferred embodiment, methods further comprise the step of extracting DNA from the homogenized sample mixture using sequence-specific nucleic acid probes. Particularly preferred are probes hybridizing to mammalian DNA.

Methods of the invention are useful to screen for the presence in a stool sample of nucleic acids indicative of colorectal cancer. Such methods comprise obtaining a representative stool sample (i.e., at least a cross-section); homogenizing the sample in a solvent having a solvent volume to stool mass ratio of at least 5:1; extracting DNA from the sample; and analyzing the DNA for characteristics of colorectal cancer. Various methods of analysis of DNA characteristics exist, such as those disclosed in co-owned, copending U.S. Patent application Serial No. 08/700,583, incorporated by reference herein.

Methods of the invention also comprise obtaining a representative (i.e., cross-sectional) sample of stool and homogenizing the stool in a buffer, such as a buffer comprising a detergent and a proteinase and optionally a DNase inhibitor.

The methods of the invention are especially and most preferably useful for detecting DNA characteristics indicative of a subpopulation of transformed cells in a representative stool sample. The DNA characteristics may be, for example, mutations, including point mutations, deletions, additions, translocations, substitutions, and loss of heterozygosity. Methods of the invention may further comprise a visual examination of the colon. Finally, surgical resection of abnormal tissue may be done in order to prevent the spread of cancerous or precancerous tissue.

Accordingly, methods of the invention provide means for screening for the presence of a cancerous or precancerous subpopulation of cells in a heterogeneous

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sample, such as a stool sample. Methods of the invention reduce morbidity and mortality associated with lesions of the colonic epithelium. Moreover, methods of the invention comprise more accurate and convenient screening methods than are currently available in the art, because such methods take advantage of the increased yield of relevant DNA. Further aspects and advantages of the invention are contained in the following detailed description thereof.

DESCRIPTION OF THE DRAWINGS

Figure 1 is a representation of a partial nucleotide sequence of the kras gene (base pairs 6282-6571) and the positions of capture probe CP1, PCR primer A1, and PCR primer B1, in relation to the kras nucleotide sequence.

Figure 2 is an image produced using a Stratagene Eagle Eye II Still Video System (Stratagene, La Jolla, CA), of the results of a gel electrophoresis run with the uncut DNA extracted as described in Example 2.

Figure 3 is an image produced using a Stratagene Eagle Eye II Still Video System (Stratagene, La Jolla, CA), of the results of a gel electrophoresis run with the DNA extracted as described in Example 3.

DETAILED DESCRIPTION OF THE INVENTION

The invention provides improved methods for extraction of nucleic acids from stool. According to methods of the invention, the yield of nucleic acids extracted from stool is increased by homogenizing the stool in a buffer at optimal ratio of buffer volume to stool mass. Improved nucleic acid yields allow nucleic acid analysis of stool samples to be conducted more efficiently with less stool volume.

In preferred methods of the invention a stool sample obtained for analysis comprises at least a cross-section of a whole stool. As provided in co-owned, copending U.S. patent application, Serial No. 08/699,678, incorporated by reference herein, cells and cellular debris from the colonic epithelium is deposited onto and into stool in a longitudinal streak. Obtaining at least a cross-section of a stool ensures that a representative sampling of colonic epithelial cells and cellular debris is analyzed.

Once the stool sample is collected, it is homogenized in a physiologically acceptable solvent. A preferred means of homogenization employs agitation with glass

beads. A preferred solvent is a physiologically-compatible buffer comprising, for example, 1M Tris, 0.5M EDTA, 5M NaCl and water to a final concentration of 500mM Tris, 16mM EDTA and 10mM NaCl at pH 9. The buffer acts as a solvent to disperse the solid stool sample during homogenization. Applicants have discovered that increasing the volume of solvent in relation to solid mass of the sample results in increased yields of DNA.

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According to methods of the invention, solvent (buffer) is added to the solid sample in a solvent volume to solid mass ratio of at least about 5:1. The solvent volume to solid mass ratio is preferably in the range of about 10:1 to about 30:1, and more preferably in the range of about 10:1 to about 20:1. Most preferably, the solvent volume to solid mass ratio is about 10:1.

In preferred methods of the invention, total DNA is isolated from stool homogenate. The homogenized mixture is centrifuged to form a pellet made up of cell debris and stool matter, and a supernatant containing nucleic acid and associated proteins, lipids, etc. The supernatant is treated with a detergent, such as 20% SDS, and enzymes capable of degrading protein (e.g., Proteinase K). The supernatant is then Phenol-Chloroform extracted. The resulting purified nucleic acids are then precipitated by means known in the art. A variety of techniques in the art can then be employed to manipulate the resulting nucleic acids, including further purification or isolation of specific nucleic acids.

In a preferred embodiment of the invention, extracted nucleic acids are placed in a physiologically compatible buffer, such as guanidine isothiocyanate (GITC). Capture probes are then added to the mixture to hybridize to target DNA in order to facilitate selective removal of target DNA from the sample.

Sequence specific capture of target DNA can be accomplished by initially denaturing sample DNA to form single-stranded DNA. Then, a sufficient quantity of sequence specific oligonucleotide probe that is complementary to at least a portion of a target polynucleotide (e.g., a sequence in or near the p53 allele) is added. The probe sequence (labeled with biotin) is allowed to hybridize to the complementary target DNA sequence. Beads coated with avidin or streptavidin are then added and attach to the

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biotinylated hybrids by affinity-binding. The beads may be magnetized to facilitate isolation.

After separation of probe-target hybrids, the resultant DNA is washed repeatedly to remove inhibitors, including those commonly introduced via the capture probe technique. In the methods of the present invention, washes are preferably carried out approximately four times with 1M GITC and 0.1% detergent, such as Igepal (Sigma). The initial washes are then preferably followed by two washes with a standard wash buffer (such as Tris-EDTA-NaCI) to remove the GITC from the mix, since GITC is a known inhibitor of DNA polymerases, including those associated with PCR.

Finally, the target DNA is eluted into a small volume of distilled water by heating. Assays using polymerase chain reaction (PCR), restriction fragment length polymorphism (RFLP) analysis or other nucleic acid analysis methods may be used to detect DNA characteristics indicative of a disorder, such as colorectal cancer or precancer. Several particularly useful analytical techniques are described in co-pending applications Serial Number 08/700,583, 08/815,576 and _______[Attorney Docket No. EXT-007], the disclosures of which are incorporated herein by reference.

Methods of the present invention are particularly useful for isolation and analysis of nucleic acids that encompass genes that have mutations implicated in colorectal cancer, such as kras. The kras gene has a length of more than 30 kbp and codes for a 189 amino acid protein characterized as a low-molecular weight GTP-binding protein. The gene acquires malignant properties by single point mutations, the most common of which occurs at the 12th amino acid. Several studies have confirmed that approximately 40% of primary colorectal adenocarcinoma cells in humans contain a mutated form of the kras gene. Accordingly, the kras gene is a particularly suitable target for the methods of colorectal cancer detection of the present invention.

Toward this end, applicants have constructed a suitable exemplary capture probe directed to the kras nucleotide sequence. The capture probe, designated CP1, has the following sequence: 5' GCC TGC TGA AAA TGA CTG AAT ATA AAC TTG TGG TAG T 3' (SEQ, ID NO: 1), and is preferably biotinylated at the 5' end in order to facilitate isolation. As illustrated more fully below, CP1 is effective in the sequence specific capture of kras DNA.

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Suitable PCR primers for the analysis of extracted kras DNA sequence have also been determined. Primer A1 has the sequence: 5' C CTG CTG AAA ATG ACT GAA 3' (SEQ ID NO: 2), and Primer B1 has the sequence: 5' CAT GAA AAT GGT CAG AGA AA 3' (SEQ ID NO: 3). The PCR primers A1 and B1, as well as capture probe CP1, are depicted in Figure 1, showing their relation to the kras nucleotide sequence, base pairs 6282-6571 (SEQ ID NO: 4). One skilled in the art can construct other suitable capture probes and PCR primers for kras or other target genes or nucleotide sequences, using techniques well known in the art.

The following examples provide further details of methods according to the invention. However, numerous additional aspects of the invention will become apparent upon consideration of the following examples.

Example 1

Stool Sample Preparation

Voided stool was collected from a patient and a cross-sectional portion of the stool was removed for use as a sample. After determining the mass of the sample, an approximately 10x volume of Tris-EDTA-NaCl lysis buffer was added to the solid sample in a test tube. The final concentration of the buffer was 500mM Tris, 16mM EDTA and 10mM NaCl, at a pH of about 9.0. Four 10mm glass balls were placed in the tube and the tube and contents were homogenized in an Exactor II shaker for 15 minutes. The homogenized mixture was then allowed to stand 5 minutes at room temperature. The tube was then centrifuged for 5 minutes at 10,000 rpm in a Sorvall Centrifuge, and the supernatant was transferred to a clean test tube. A 20% SDS solution was added to the tube to a final concentration of 0.5%. Proteinase K was also added to the tube to a final concentration of 500mg/ml. The tube was then incubated overnight at 37°C.

After incubation, the contents of the tube were extracted with an equal volume of phenol/chloroform and centrifuged at 3500 rpm for 3 minutes. The aqueous layer was then transferred to a new tube and extracted three (3) times with equal volumes of chloroform and centrifuged at 3500 rpm for 3 minutes. The aqueous layer was then transferred to a new tube and 0.1x volume of 3M NaOAc was added to the aqueous portion, which was then extracted with an equal volume of isopropanol, and centrifuged

for 5 minutes at 12,000 rpm. The supernatant was discarded, and the pellet was washed with 10ml of 70% ethanol, and centrifuged at 12,000 rpm for 5 minutes. The supernatant was discarded and the pellet containing isolated DNA was dried by inverting the tube.

5 Example 2

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A comparative analysis of solvent volume to mass ratios was conducted. Three separate stool samples were prepared as described above. A first sample, designated SS88-3x, was homogenized in buffer at a volume to mass ratio of 3:1. A second sample, designated SS88-5x, was homogenized at a ratio of 5:1; and a third sample, designated SS88-10x, was homogenized at a ratio of 10:1.

Total DNA from each sample was resuspended in 100 ul of 100 mM Tris, 10 mM EDTA buffer and 10 ul aliquots were loaded onto a 4% agarose gel for electrophoresis at 125 V constant voltage for about one hour. The results are shown in Figure 2. As shown in Figure 2, the yield of total DNA increased as the ratio of solvent to mass increased from 3x to 10x.

Example 3

A second set of four equivalent samples was prepared from a single stool sample. Each of the four samples was of equal mass, and was homogenized as described in Example 1 at a solvent volume to stool mass ratio of 5:1, 10:1, 20:1, and 30:1, respectively. After homogenization each sample was subdivided into 8 aliquots, 4 treated with RNase, and 4 untrreated. Total DNA was then isolated as described above and analyzed on agarose gels.

The results are shown in Figure 3. As shown, a ratio of 10:1 produced the greatest yield of nucleic acids. Figure 3 also shows the effect of RNase treatment on the yield of DNA from each stool sample. As shown in the Figure, RNase treatment virtually eliminates RNA from the sample, but leaves DNA intact. The results indicate that optimal solvent volume to stool mass ratios greatly increase DNA yield from stool samples.

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Example 4

Sequence-Specific Capture of target DNA.

Once extracted from stool, specific nucleic acids are isolated using sequence-specific capture probes. Total DNA was extracted from a stool sample according to the methods described in Example 1. The pelletized DNA was resuspended in 1ml of TE buffer. A 100 µl aliquot of this solution was removed to a new tube and 100 µl of 6M guanidine isothiocyanate (GITC) was added to a final concentration of 3M GITC. A vast excess of biotinylated kras capture probe CP1 was the added to the sample. The mixture was heated to 95°C for 5 minutes to denature the DNA, then cooled to 37°C for 5 minutes. Finally, probe and target DNA were allowed to hybridize for 30 minutes at room temperature. Streptavidin-coated magnetized beads (320 mg) (Dynal Corp.) were suspended in 400 µl distilled water and added to the mixture. After briefly mixing, the tube was maintained at room temperature for 30 minutes.

Once the affinity binding was completed, a magnetic field was applied to the sample to draw the magnetized isolation beads (both with and without hybridized complex out of the sample. The beads were then washed four (4) times in 1M GITC/0.1% Igepal (Sigma, St. Louis, MO) solution for 15 minutes, followed by two (2) washes with wash buffer (TE with 1M NaCl) for 15 minutes in order to isolate complexed streptavidin. Finally, 10 µl distilled water was added to the beads and heated at 95°C for 3 minutes to elute the DNA. Sequencing and/or gel electrophoresis enable confirmation of the capture of kras-specific DNA.

Accordingly, methods of the invention produce increased yields of DNA from stool, thereby allowing more efficient sequence-specific capture of target nucleic acid. Methods of the invention provide improvements in the ability to detect disease-related nucleic acid mutations present in stool. The skilled artisan will find additional applications and embodiments of the invention useful upon inspection of the foregoing description of the invention. Therefore, the invention is limited only by the scope of the appended claims.

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SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: SHUBER, Anthony P. LAPIDUS, Stanley N. RADCLIFFE, Gail E.
 - (ii) TITLE OF INVENTION: Methods for Stool Sample Preparation
 - (iii) NUMBER OF SEQUENCES: 4
- ____(iv) CORRESPONDENCE_ADDRESS:
 - (A) ADDRESSEE: Testa, Hurwitz & Thibeault
 - (B) STREET: 125 High St.
 - (C) CITY: Boston
 - (D) STATE: MA
 - (E) COUNTRY: USA
 - (P) ZIP: 02110
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk

 - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:

 - (A) NAME: MEYERS, Thomas C. (B) REGISTRATION NUMBER: 36,989
 - (C) REFERENCE/DOCKET NUMBER: EXT-006PC
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (617) 248-7000 (B) TELEFAX: (617) 248-7100
 - (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 37 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

GCCTGCTGAA AATGACTGAA TATAAACTTG TGGTAGT

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- (2) INFORMATION FOR SEQ ID NO:2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 19 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
CCTGCTGAAA ATGACTGAA	19
(2) INFORMATION FOR SEQ ID NO:3:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
CATGAAAATG GTCAGAGAAA	20
(2) INFORMATION FOR SEQ ID NO:4:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 307 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
GTACTGGTGG AGTATTGAT AGTGTATTAA CCTTATGTGT GACATGTTCT AATATAGTCA	60
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TTGGAGCTGG TGGCGTAGGC AAGAGTGCCT TGACGATACA GCTAATTCAG AATCATTTTG	180
TGGACGAATA TGATCCAACA ATAGAGGTAA ATCTTGTTTT AATATGCATA TTACTGGTGC	240
AGGACCATTC TITGATACAG ATAAAGGITT CTCTGACCAT TITCATGTAC AGAAGTCCTT	300
GCTAAGA	30

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CLAIMS

What is claimed is:

1	1.	A method for extracting DNA from stool, comprising:
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- 2 homogenizing a stool sample in a solvent for DNA in order to form a
- 3 homogenized sample mixture having a solvent volume to stool mass ratio of at
- 4 least 5:1; and
- 5 extracting DNA from said homogenized sample mixture.
- 1 2. The method of claim 1 wherein the solvent volume to stool mass ratio is from
- 2 about 10:1 to about 30:1.
- 1 3. The method of claim 2 wherein the solvent volume to stool mass ratio is about
- 2 10:1 to about 20:1.
- 1 4. The method of claim 2 wherein the solvent volume to stool mass ratio is about
- 2 10:1.
- 1 5. The method of claim 1 wherein the solvent comprises a physiologically
- 2 compatible buffer.
- 1 6. The method of claim 5 wherein the buffer comprises Tris-EDTA-NaCl.
- 1 7. The method of claim 6 wherein the Tris-EDTA-NaCl buffer comprises a final
- 2 concentration of about 50mM Tris, about 16 mM EDTA and about 10mM NaCl at
- 3 about pH 9.0.
- 1 8. The method of claim 1 wherein the solvent comprises guanidine isothiocyanate
- 2 buffer.
- 1 9. The method of claim 8 wherein the guanidine isothiocyanate buffer comprises a
- 2 final concentration of from about 1 to about 5 M.

1 2	10.	The method of claim 9 wherein the guanidine isothiocyanate buffer comprises a final concentration of about 3 M.
1 2	11.	The method of claim 1 wherein said extracting step further comprises contacting said DNA with a sequence-specific capture probe.
1	12.	The method of claim 1 wherein said solvent comprises a detergent and a proteinase.
1	13.	The method of claim 1 wherein said DNA is human DNA.
1	14.	A method of screening for the presence of a colorectal cancerous or pre- cancerous lesion in a patient, the method comprising the steps of:
3		obtaining a sample comprising at least a cross-sectional portion of a stool voided by the patient;
5 6		homogenizing the sample in a solvent in order to form a homogenized sample mixture having a solvent volume to stool mass ratio of at least 5:1;
7		extracting target human DNA from said homogenized sample mixture; and
8 9		analyzing the extracted target human DNA for DNA characteristics indicative of the presence of said colorectal cancerous or pre-cancerous lesion.
1 2	15.	The method of claim 14 wherein said analyzing step comprises amplifying the DNA with a polymerase chain reaction.
1 2	16.	The method of claim 14 wherein said DNA characteristics comprise a loss of heterozygosity encompassing a polymorphic locus.
1	17.	The method of claim 14 wherein said DNA characteristic is a mutation.
1	18.	The method of claim 17 wherein said mutation is selected from the group consisting of loss of heterozygosity and microsatellite instability.

- 1 19. The method of claim 14 wherein said DNA characteristics comprise a deletion in a tumor suppressor allele.
- The method of claim 14 wherein said analyzing step comprises determining
 whether a difference exists in said sample between a number X of a first allele
 known or suspected to be mutated in a subpopulation of cells in the sample and
 a number Y of a second allele that is known or suspected not to be mutated in a
 subpopulation of cells in the sample, the presence of a statistically-significant
 difference being indicative of a mutation in a subpopulation of cells in the
 sample and the potential presence of a cancerous or precancerous lesion.
- The method of claim 14 wherein said analyzing step comprises determining
 whether a difference exists between a number of a target tumor suppressor
 allele in the sample and a number of a non-cancer-associated reference allele in
 the sample, the presence of a statistically-significant difference being indicative
 of a deletion of the target tumor suppressor allele in a subpopulation of cells in
 the sample and the potential presence of a cancerous or precancerous lesion.
- The method of claim 14 wherein said analyzing step further comprises the steps of:
- a) detecting an amount of a maternal allele at a polymorphic locus in the sample;
- b) detecting an amount of a paternal allele at the polymorphic locus in the
 sample; and
- 7 c) determining whether a difference exists between the amounts of maternal and paternal allele,
- the presence of a statistically-significant difference being indicative of a deletion at the polymorphic locus in a subpopulation of cells in the sample and the potential presence of a lesion.

1	23.	The method	of claim 22 wherein said polymorphic locus is a single	base
2	polym	norphism and	is heterozygous between said maternal and paternal al	leles.
1	24.	The method	of claim 22 wherein said detecting steps comprise,	
2		a)	hybridizing probe to a portion of said polymorphic local	us on both
3			maternal and paternal alleles that is immediately adja	cent to said
4			single-base polymorphism;	
5		b)	exposing said sample to a mixture of detectably-label	ed dideoxy
6			nucleoside triphosphates under conditions which allo	w appropriate
7			binding of said dideoxy nucleoside triphosphates to s	aid single-
8			base polymorphism;	
9	•	c)	washing the sample; and	
10		d)	counting an amount of each detectably-labeled dideo	xy nucleoside
11			triphosphate remaining for the sample.	
1	25.	The method	d of claim 24 wherein said detectable label is selected f	rom the group
2	cons	sisting of radio	oisotopes, fluorescent compounds, and particles.	
1	26.	The metho	d of claim 14 wherein said analyzing step comprises a r	nethod for
2	dete	cting heteroz	ygosity at a single-nucleotide polymorphic locus, compr	ising the steps
3	of:			
4		a)	hybridizing probes to a sequence immediately adjace	ent to a single
5			base polymorphism;	
6		. b)	exposing the sample to a plurality of different labele	d dideoxy
7	٠		nucleotides	
8		c)	washing the sample;	
9		d)	determining which of said dideoxy nucleotides are in	ncorporated
10			into said probes; and	

11	e) detecting heterozygosity at the single-nucleotide polymorphic site
12	as the detection of two dideoxy nucleotides having been
13	incorporated into the probe.
1	27. The method of claim 14 wherein said analyzing step comprises:
2	(a) exposing the sample to a plurality of a first oligonucleotide probe
3	and to a plurality of a second oligonucleotide probe under hybridization conditions,
4	thereby to hybridize
5	(1) said first oligonucleotide probes to copies of a first
6	polynucleotide segment characteristic of wild-type cells of the organism, and
7	(2) said second oligonucleotide probes to copies of a second
. ' 8	polynucleotide segment characteristic of a wild-type genomic region suspected to be
	deleted or mutated in colorectal cancer cells;
9	deleted of flutated in colorectal carries with,
10	(b) detecting a first number of duplexes formed between said first
11	probe and said first segment and a second number of duplexes formed between said
12	second probe and said second segment; and
	(c) determining whether there is a difference between the number of
13	
14	duplexes formed between said first probe and said first segment and the number of
15	duplexes formed between said second probe and said second segment,
16	the presence of a statistically-significant difference being indicative of the
17	presence in said sample of a colorectal cancer or precancerous lesion.
.,	
1	28. The method of claim 27 wherein said first and second oligonucleotide probes
2	each are coupled to a distinct detectable label.
1	29. The method of claim 27 wherein
2	said first oligonucleotide probes are attached to a first particle in a ratio of
3	one first oligonucleotide probe to one particle and said second oligonucleotide probes

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4	are attached to a second particle detectably distinct from said first particle in a ratio of
5	one second oligonucleotide probe to one second particle, wherein
6 7 8 9	said detecting step comprises separating hybridized from unhybridized first and second oligonucleotide probes and subsequently passing hybridized first and second oligonucleotide probes through a detector to determine said first and second numbers.
•	
1	30. The method of claim 29 wherein said first and second particles are of detectably
2	different sizes.
1	31. The method of claim 29 wherein said first and second particles are of detectably
2	different colors.
1	32. The method of claim 27 further comprising, prior to step a) the steps of
2	converting double-stranded DNA in said sample to single-stranded DNA and removing
3	complement to said first and second polynucleotide segments.
1 2	33. The method of claim 32 wherein said removing step comprises hybridizing said complement to a nucleic acid probe attached to a magnetic particle and subsequently
3	removing said magnetic particle from the sample.
1	34. The method of claim 14 wherein said analyzing step comprises a method for
2	detecting a nucleic acid sequence change in a target allele in the sample, comprising
3	the steps of:
4	(a) determining
5	(i) an amount of wild-type target allele in the sample, and
6	(ii) an amount of a reference allele in the sample; and
7	(b) detecting a nucleic acid sequence change in the target allele in the
8	sample,

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9		a statistically significant difference in the amount wild-type target allele
0		and the amount of reference allele obtained in said determining step being
1		indicative of a nucleic acid sequence change.
1	35 .	The method according to claim 34 wherein said determining step comprises
2	ехро	sing said sample to a first oligonucleotide probe capable of hybridizing with a
3	porti	on of said wild-type allele and to a second oligonucleotide probe capable of

hybridizing to a portion of said reference allele, and removing from said sample any unhybridized first or second oligonucleotide probe. 5

Fig. 1

5'-GT ACT GGT GGA GTA TTT GAT AGT GTA TTA ACC TTA

TGT GTG ACA TGT TCT AAT ATA GTC ACA TTT TCA TTA TTT

TTA TTA TAA GGC CTG CTG AAA ATG ACT GAA TAT AAA

A1

CTT GTG GTA GTT GGA GCT GGT GGC GTA GGC AAG AGT GCC

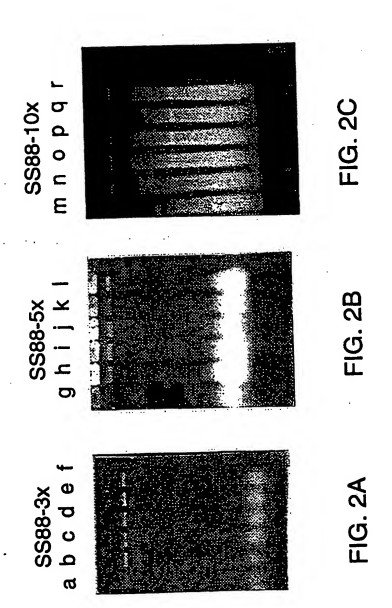
TTG ACG ATA CAG CTA ATT CAG AAT CAT TTT GTG GAC GAA

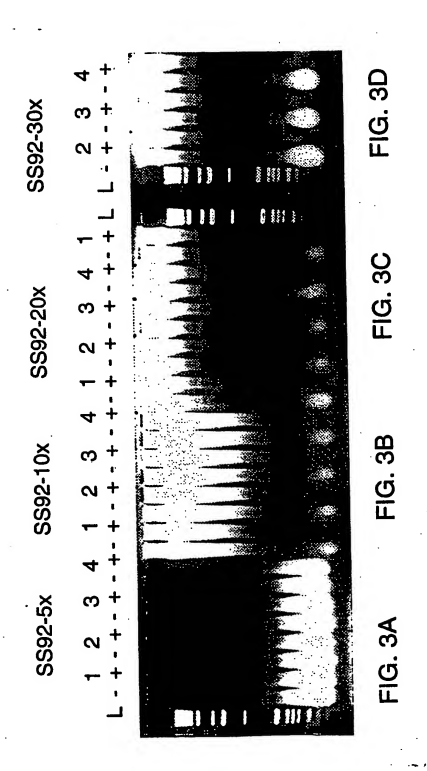
TAT GAT CCA ACA ATA GAG GTA AAT CTT GTT TTA ATA

TGC ATA TTA CTG GTG CAG GAC CAT TCT TTG ATA CAG ATA

AAG GTT TCT CTG ACC ATT TTC ATG TAC AGA AGT CCT TGC

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A. CLASSI IPC 6	FICATION OF SUBJECT MATTER C12Q1/68			
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B. FIELOS	SEARCHED			
Minimum do IPC 6	cumentation searched (classification system followed by classification C 120	n symbols)		
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Electronic d	ata base consulted during the international search (name of data bas	e and, where practical.	search terms used)	
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